Direct Reconstruction of Kinetic Parameter Images from Dynamic PET Data

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Abstract—Our goal in this paper is the estimation of kinetic model parameters for each voxel corresponding to a dense 3D PET image. Typically, the activity images are first reconstructed from PET sinogram frames at each measurement time, and then the kinetic parameters are estimated by fitting a model to the reconstructed time-activity response of each voxel. However, this “indirect” approach to kinetic parameter estimation tends to reduce signal-to-noise ratio (SNR) because of the requirement that the sinogram data be divided into individual time frames.

In 1985, Carson and Lange proposed [1], but did not implement, a method based on the expectation-maximization (EM) algorithm for direct parametric reconstruction. The approach is “direct” because it estimates the optimal kinetic parameters directly from the sinogram data, without an intermediate reconstruction step. However, direct voxel-wise parametric reconstruction remained a challenge due to the unsolved complexities of inversion and spatial regularization.

In this work, we demonstrate and evaluate a new and efficient method for direct voxel-wise reconstruction of kinetic parameter images using all frames of the PET data. The direct parametric image reconstruction is formulated in a Bayesian framework, and uses the parametric iterative coordinate descent (PICD) algorithm to solve the resulting optimization problem [2]. The PICD algorithm is computationally efficient and is implemented with spatial regularization in the domain of the physiologically relevant parameters. Our experimental simulations of a rat head imaged in a working small animal scanner indicate that direct parametric reconstruction can substantially reduce root mean square error (RMSE) in the estimation of kinetic parameters, as compared to indirect methods, without appreciably increasing computation.

Index Terms—tomography, iterative reconstruction, dynamic PET, kinetic modeling, regularization

I. INTRODUCTION

Positron Emission Tomography (PET) is a powerful molecular imaging technique with the sensitivity to detect picomolar quantities of a labelled tracer with reasonable (seconds to minutes) temporal resolution. Through the application of kinetic models, the dynamic PET data can be transformed into physiological parameters that indicate the functional state of the imaged tissue. Ideally, one would like to reconstruct parametric images from PET data i.e., images which specify the estimated kinetic parameters for each voxel in the imaged volume. Such parametric images could serve many uses. For example, they may be particularly desirable when testing a new tracer whose sites of action are not completely known. In the brain, parametric images might be useful in identifying new brain circuits or discovering unsuspected connectivity between disparate brain regions. As new tracers continue to be developed with greater specificity to nonspecific binding ratios, the impetus grows for looking at their uptake in all regions of the brain, rather than in a few pre-selected regions of interest.

For example, $^{18}$F-fallypride, a high affinity dopamine tracer, can be used to image dopamine receptors outside the striatum because the non-specific background is low [3].

This paper introduces a novel algorithm for directly reconstructing parametric images from PET sinogram data. We demonstrate that this method can generate parametric images with superior quality; and, perhaps surprisingly, we also show that it has computational requirements that are similar to a two-step approach of iterative reconstruction followed by kinetic parameter estimation.

Kinetic compartmental models are often used to describe the movement of a tracer between different physically or chemically distinct states or compartments [4]–[6]. The exchange of tracer between these compartments can be modeled by a system of first order ordinary differential equations (ODEs) whose coefficients are the kinetic parameters. The resulting kinetic models have been validated as producing reliable quantitative indices of various clinically and scientifically important physiological processes [7]–[17].

In some cases, a single set of kinetic parameters can describe the tracer behavior in a homogeneous region of tissue such as the myocardium or perhaps the entire striatum in brain images. If the region of interest can be delineated using some form of segmentation, then the PET activity can be averaged over the region at each time frame and a single set of kinetic parameters can be estimated by fitting a single kinetic model to the time sequence of average activities. This case is illustrated in Fig. 1. The PET data are first reconstructed into $K$ time frames, then a region of interest (ROI) is segmented from each frame, and a single set of kinetic parameters is fit to the regional-average time sequence. These ROI-based methods may be further classified into linear methods and nonlinear methods. Linear techniques [18]–[23] transform the data, so that the parameters of interest can be estimated by linear regression methods, while nonlinear techniques [5], [6] generally estimate the kinetic parameters by iteratively minimizing a properly weighted distance metric between the reconstructed time-activity curves and the model output.
Recently, there has been increasing interest in the formation of parametric images which model the kinetic behavior of each voxel individually. This approach is more appropriate when the volume cannot be effectively segmented into homogeneous regions that would be modeled with a single kinetic parameter set. Existing approaches to the creation of parametric images can be roughly categorized as “indirect”, “semi-direct”, and (our new method) “direct” [24] reconstruction.

Indirect methods work by first reconstructing the PET emission images for each of the $K$ measurement times, and then estimating the kinetic parameters at each voxel. The primary difficulty of the indirect approach is that the low signal-to-noise ratio of the time-activity curve (TAC) for each voxel makes accurate estimation of parameters difficult. To improve estimation accuracy, O’Sullivan et al. [25] applied ridge regression techniques to regularize the parameters using prior knowledge of their means and variances derived from the analysis of a reasonably large patient group. Huang et al. [26] applied a spatial smoothing step between the iterations of a nonlinear estimation process at each voxel. Later, Zhou et al. [27], [28] developed a two-stage algorithm whereby the kinetic parameters were estimated first using standard nonlinear techniques. In a second step the initial results were smoothed spatially and used to constrain the final estimates. (This method is further discussed in section IV-B.) Kimura et al. [29] and Zhou et al. [30] have developed algorithms that cluster the images before estimation and regularize the data within the clusters.

Semi-direct algorithms, as they are sometimes named, attempt to improve signal-to-noise by constraining the possible choices of time-courses for each voxel via signal subspaces or splines. Kao et al. [31] and Narayanan et al. [32] used principal component analysis (PCA) to decorrelate the sinograms in time. Similarly, Wernick et al. [33] applied PCA decomposition of PET data followed by reconstruction of tracer concentrations in the principal-component space. Nichols et al. [34], and Reutter et al. [35], [36] proposed reconstruction methods that use a b-spline specification of the time-activity curves. Kinetic parameters must then be estimated from the b-spline representation. It is important to note that spline-based methods have certain computational advantages when processing list mode data. However, the two-step process of first computing spline coefficients and then kinetic parameters still results in a loss of optimality, particularly if the number of spline coefficients is much larger than the number of kinetic parameters.

Ideally, one would like to estimate directly the space-domain kinetic parameters from the measured sinogram data. In fact, Carson and Lange [1] proposed direct estimation of kinetic parameters from PET data in 1985. In that paper, the authors outlined a general framework for a direct reconstruction algorithm based on expectation-maximization (EM) [37] iterations. Unfortunately, the Carson and Lange direct parametric reconstruction algorithm has never, to our knowledge, been fully implemented for nonlinear estimation of a dense set of voxels. Limber et al. [38] proposed an algorithm for direct parametric reconstruction using maximum likelihood (ML) estimation of kinetic parameters from PET data, but only demonstrated the algorithm for an $8 \times 8$ array of voxels. A number of authors have implemented direct nonlinear parameter estimation methods that were designed for segmented ROI’s [39]–[44] rather than a dense set of voxels. In an alternative approach, Meikle et al. [45] first pre-computed the time-activity curves for a range of possible nonlinear parameters and then directly reconstructed the linear weights for each of the nonlinear “basis” curves. Similarly, Matthews et al. [46] obtained pre-defined time-curves from other patients’ reconstructions and used the EM algorithm to compute the weights of each curve. In other work, Carson [47] proposed an ML framework to estimate the ROI values from the projections, and Farncombe et al. [48] estimated organ uptake parameters that were incorporated into the reconstruction algorithm for dSPECT applications.

In this paper, we present an algorithm for direct nonlinear estimation of space-domain kinetic parameters in a dense volume of voxels. Our novel parametric reconstruction algorithm, which we call parametric iterative coordinate descent (PICD) [2], is in the spirit of Carson and Lange’s method. However, PICD is a completely specified and implemented algorithm (See Appendix A) which we show to be computationally efficient with robust convergence properties. In fact, the computation required for parametric reconstruction using PICD is comparable to that required for more conventional maximum $a$ posteriori (MAP) reconstruction of an image sequence from PET sinogram data. In other words, it is our claim that direct parametric reconstruction can have comparable computational requirements to indirect methods (recall that indirect methods require an initial reconstruction of all the data). The key to computational efficiency of the PICD method is the use of state variables and nested optimization to decouple the nonlinearities of the forward tomographic model, the nonlinear kinetic model, and the Bayesian prior model. Notably, PICD is designed to compute the MAP estimate of the kinetic parameters using a prior distribution defined on any well-behaved transformation of the parameter space. This allows the regularization to be applied to the parameters that are deemed to be physiologically important. Simulation results, presented below, indicate that the PICD-generated parametric reconstructions have lower mean squared error and better visual quality than the best indirect methods.

Section II reviews the 2-tissue compartment model and the set of ODE’s that govern a tracer’s kinetics. Section III introduces the PICD algorithm for direct parametric reconstruction and gives a detailed description of its implementation. Section IV first reviews some existing methods for image domain parameter estimation, and then suggests a useful method for regularization of pixel-wise approaches. Section V compares the computational complexity of the proposed methods. Section VI presents simulation results. Discussion and Conclusion follow the results.
II. 2-TISSUE COMPARTMENT MODEL

In this paper, we used a 2-tissue compartment model to describe the kinetic processes that are represented by the signal from each voxel of a reconstructed image. This model is commonly used to describe the uptake and retention of an analog of glucose, 2-deoxy-2-[\textsuperscript{18}F]fluoro-D-glucose (FDG). The model can also be properly applied to receptor ligand studies provided that there is no non-specific binding and that the tracer has been administered at sufficiently high specific activity. Figure 2 illustrates the model: \( C_P \) (pmol/ml) is the molar concentration of tracer in the plasma, \( C_F \) (pmol/ml) is the molar concentration of unbound tracer, and \( C_B \) (pmol/ml) is the molar concentration of metabolized or bound tracer. The model depends on the kinetic parameters, \( k_1, k_2, k_3, \) and \( k_4 \), which specify the tracer exchange rates between compartments in units of inverse minutes. The parameters \( k_1, k_2, \) and \( k_4 \) are first order rate constants, and \( k_3 \) is an apparent first order rate constant describing a process (metabolism or receptor-binding) that proceeds in proportion to the concentration of the labelled tracer only, as long as the number of sites available for binding do not become rate-limiting.

In addition to the above-stated parameters, there are two compound parameter groups that have ready physiological interpretations and practical application, particularly for receptor-ligand imaging: binding potential \((BP)\), and total volume of distribution \((VD)\). \( BP \) is proportional to the number of receptors and \( VD \) represents the steady state distribution of tracer between the plasma and tissue. \( BP \) and \( VD \) can be expressed in terms of the aforementioned kinetic parameters,

\[
BP = \frac{k_3}{k_4} \\
VD = \frac{k_1}{k_2} \left(1 + \frac{k_3}{k_4}\right). \tag{1}
\]

In applying the model in Fig. 2 to all voxels, we assume that the delivery of tracer is the same to all regions being imaged. In other words, the value of \( C_P \) is not a function of voxel position. However, the values of the kinetic parameters will be allowed to vary for each voxel location, \( s \). Using these assumptions, the time variation of the concentrations for a single voxel are governed by the following ordinary differential equations (ODE).

\[
\frac{dC_P(s,t)}{dt} = k_{1s}C_P(t) - (k_{2s} + k_{3s})C_F(s,t) + k_{4s}C_B(s,t) \tag{3}
\]

\[
\frac{dC_B(s,t)}{dt} = k_{3s}C_F(s,t) - k_{4s}C_B(s,t). \tag{4}
\]

In this work, \( C_P(t) \) is assumed known. In practice, it can be measured directly from arterial plasma samples during the imaging procedure \cite{6}, or it may be estimated from imaged volumes that consist primarily of blood \cite{49}--\cite{53}. The solution to the ODE’s in (3,4) is given by

\[
C_P(s,t) = \left\{ \frac{k_{1s}}{\alpha_2 - \alpha_1}[(k_{4s} - \alpha_1)e^{-\alpha_1t} + (\alpha_2 - k_{4s})e^{-\alpha_2t}]u(t) \right\} * C_P(t) \tag{5}
\]

\[
C_B(s,t) = \left\{ \frac{k_{1s}k_{3s}}{\alpha_2 - \alpha_1}e^{-\alpha_1t} - e^{-\alpha_2t}]u(t) \right\} * C_P(t) \tag{6}
\]

where \( * \) indicates continuous-time convolution, \( \alpha_1 \) and \( \alpha_2 \) are real valued constants that result from the subtraction and addition of terms in (7) respectively, and \( u(t) \) is the unit step function.

\[
\alpha_1, \alpha_2 = \frac{(k_{2s} + k_{3s} + k_{4s})}{2} + \sqrt{\left(k_{2s} + k_{3s} + k_{4s}\right)^2 - 4k_{2s}k_{4s}} \tag{7}
\]

Next, we transform the kinetic parameters \((k_1, k_2, k_3, k_4)\) to form the new parameters \((a, b, c, d)\) as shown in Table I. This transformation is important because while the parameters \((a, b, c, d)\) are well suited for optimization, \((k_1, k_2, k_3, k_4)\) are more physiologically relevant. We use \( \varphi_s = [a_s, b_s, c_s, d_s] \) to denote the parameter vector for each voxel \( s \).

The total activity concentration (e.g., in nCi/ml) for voxel \( s \) at time \( t \) is denoted by

\[
f(\varphi_s, t) = (1 - V_B) \left[ C_P(s,t) + C_B(s,t) \right] S_A e^{-\lambda t} + V_B C_{WB}(t) \\
= (1 - V_B) \left[ (a_s e^{-c_s t} + b_s e^{-d_s t})u(t) * C_P(t) \right] S_A e^{-\lambda t} + V_B C_{WB}(t) \tag{8}
\]
where \( S_A \) is the initial specific activity of the tracer (nCi/μmol), \( \lambda \) is the decay rate of the isotope (min^{-1}), \( V_B \) is a known constant for the volume fraction of the voxel that contains blood, and \( C_{WB} \) (nCi/ml) is the tracer activity concentration in whole blood (i.e., plasma plus blood cells plus other particulate matter). \(^1\) We can simplify the expression for \( f(\varphi, t) \) by defining the following functions:

\[
\begin{align*}
\alpha(c, t) & \equiv \{C_P(t) * [e^{-c_{s} t}u(t)]\} (1 - V_B) S_A e^{-\lambda t} \\
\beta(d, t) & \equiv \{C_P(t) * [e^{-d_{s} t}u(t)]\} (1 - V_B) S_A e^{-\lambda t} \\
\gamma(t) & \equiv V_B C_{WB}(t) 
\end{align*}
\]

With these definitions, \( f(\varphi, t) \) can be written as

\[
f(\varphi, t) = [a_s, b_s] \left[ \begin{array}{c} \alpha(c_s, t) \\ \beta(d_s, t) \end{array} \right] + \gamma(t) .
\]

We next define some vector and matrix notation that will be useful in discretization of the problem. \(^2\) Let \( t_0, \ldots, t_{K-1} \) be the \( K \) discrete times at which the tissue is imaged. Then we may construct the vectors:

\[
\begin{align*}
\alpha(c, t) & \equiv \{\alpha(c_s, t_0), \alpha(c_s, t_1), \ldots, \alpha(c_s, t_{K-1})\} \\
\beta(d, t) & \equiv \{\beta(d_s, t_0), \beta(d_s, t_1), \ldots, \beta(d_s, t_{K-1})\} \\
\gamma & \equiv \{\gamma(t_0), \gamma(t_1), \ldots, \gamma(t_{K-1})\}
\end{align*}
\]

Using this notation, the activity at each time for voxel \( s \) is given by the \( 1 \times K \) row vector

\[
f(\varphi, t) = [f(\varphi, t_0), f(\varphi, t_1), \ldots, f(\varphi, t_{K-1})]
\]

\[
= [a_s, b_s] \left[ \begin{array}{c} \alpha(c_s) \\ \beta(d_s) \end{array} \right] + \gamma .
\]

Let the \( N \) voxels be indexed by the values \( s = 0, 1, \ldots, N - 1 \), and let \( \varphi = \{\varphi_0, \varphi_1, \ldots, \varphi_{N-1}\} \) denote the \( 4 \times N \) matrix of parameters at all voxels. With this, we define the \( N \times K \) function

\[
F(\varphi) = \left[ \begin{array}{c} f(\varphi_0) \\ \vdots \\ f(\varphi_{N-1}) \end{array} \right]
\]

\(^1\) Notice that both \( f(\varphi, t) \) and \( C_{WB}(t) \) include the attenuation due to decay. Therefore, the sinogram data should not be decay corrected for the implementation of this method.

\(^2\) This discretization approach is equivalent to interpreting the measured concentrations as representing instantaneous concentrations measured at mid-frame.

### III. PARAMETRIC RECONSTRUCTION FROM SINOGRAM DATA

In this section, we describe our method for directly reconstructing the parametric image, \( \varphi \), from sinogram data. We will do this by first formulating a conventional scanner model under the assumption that the sinogram measurements are Poisson random variables. We will then use the kinetic model of Section II as the input to the scanner model as shown in Fig. 3. Once the complete forward model is formulated, we will present an iterative algorithm for computing the maximum a posteriori (MAP) estimate of the parametric image \( \hat{\varphi} \) from the sinogram data. Once \( \hat{\varphi} \) is computed, the activity images can be computed at any time \( t \) simply by evaluating \( F(\varphi, t) \) using the kinetic model equations of (8).

#### A. Scanner Model

Let \( Y_{mk} \) denote the sinogram measurement for projection \( 0 \leq m < M \) and time frame \( 0 \leq k < K \), and let \( Y \) be the \( M \times K \) matrix of independent Poisson random variables that form the sinogram measurements. Furthermore, let \( A \) be the forward projection matrix, with elements \( A_{ms} \) (counts/mlCi), and let \( \mu \) be the number of accidental coincidences. Then the expected number of counts for each measurement at a given time, \( t_k \) is given by

\[
E[Y_{mk}|F(\varphi, t_k)] = \sum_{s=0}^{N-1} A_{ms} f(\varphi_s, t_k) + \mu .
\]

This relationship can be compactly expressed using matrix notation as

\[
E[Y|F(\varphi)] = AF(\varphi) + \mu .
\]

It is easily shown that under these assumptions the probability density for the sinogram matrix is given by [54]

\[
p(Y|\varphi) = \frac{1}{\prod_{k=0}^{K-1} \prod_{m=0}^{M-1} \left( A_{ms} F(\varphi, t_k) + \mu \right) Y_{mk}!} .
\]

where \( A_{ms} \) is the \( mth \) row of the system matrix, \( A \). The log likelihood of the sinogram matrix is then given by

\[
LL(Y|\varphi) = \sum_{k=0}^{K-1} \sum_{m=0}^{M-1} Y_{mk} \log(A_{ms} F(\varphi, t_k) + \mu) - (A_{ms} F(\varphi, t_k) + \mu) - \log(Y_{mk}) .
\]

This is a very general formulation. For specific scanners, the form of the system matrix \( A \) may vary considerably, and accurate determination of the matrix \( A \) can be critical to obtaining accurate tomographic reconstructions [55],[56].
that minimizes this cost function. The MAP reconstruction, \( \hat{\varphi} \), is similar to the ICD algorithm used in conventional PET metric iterative coordinate descent (PICD). This algorithm is computed efficiently by an algorithm which we call parametric image reconstruction \([54]\), but it is adapted to account for the nonlinear parameters of the compartmental model. PICD sequentially updates the parameters of each voxel thereby monotonically decreasing the cost function given in Equation (22). When \( F(\varphi) \) is a nonlinear function, the PICD algorithm reduces computation by decoupling the dependencies between the compartment model nonlinearities and the forward tomography model.

In order to compute a PICD voxel update, we must compute

\[
\varphi_s \leftarrow \arg \min_{\varphi_s} C(Y|\varphi_s) .
\]

To do this efficiently, we use the second order Taylor expansion of the change in the cost function.

Suppose we are updating the parameters of voxel \( s \) from \( \varphi_s = [a_s, b_s, c_s, d_s]^T \) to \( \tilde{\varphi}_s = [a_s, b_s, c_s, d_s]^T \), and that we represent the change in the time response function of voxel \( s \) by the \( 1 \times K \) vector function

\[
\Delta f(\tilde{\varphi}_s, \varphi_s) = f(\tilde{\varphi}_s) - f(\varphi_s) .
\]

We next define a simplified cost functional

\[
\Delta C(\tilde{\varphi}_s, \varphi_s) = -LL(Y|\tilde{\varphi}_s) + LL(Y|\varphi_s) + \sum_{r \in \partial s} g_{s-r} || T(\tilde{\varphi}_s) - T(\varphi_r) ||_W^2 .
\]

Notice that since \( \Delta C(\tilde{\varphi}_s, \varphi_s) \) is equal to the change in the cost functional \( C(Y|\tilde{\varphi}_s) \) within a constant, so it may be used to compute the voxel update of (25). The value of \( \Delta C(\tilde{\varphi}_s, \varphi_s) \) can then be locally approximated with a second order Taylor series as

\[
\Delta C(\tilde{\varphi}_s, \varphi_s) \approx \Delta f(\tilde{\varphi}_s, \varphi_s) \theta_1 + \frac{1}{2} || \Delta f(\tilde{\varphi}_s, \varphi_s) ||_{\theta_2}^2 + \sum_{r \in \partial s} g_{s-r} || T(\tilde{\varphi}_s) - T(\varphi_r) ||_W^2 .
\]

where \( \partial s \) denotes the set of voxels that are 8-neighbors of voxel \( s \), \( \theta_1 \) is a \( K \times 1 \) vector, \( \theta_2 \) is a \( K \times K \) diagonal matrix, and \( || x ||_{\theta_2}^2 = x^T \theta_2 x \). Here the values of \( \theta_1 \) and \( \theta_2 \) consist of the first and second derivatives respectively of the log likelihood function evaluated at each time frame. These derivatives at time frame \( k \) can be iteratively updated using the equations of the conventional iterative coordinate descent (ICD) algorithm \([54]\), given in (28) and (29).

\[
[\theta_1]_k \leftarrow \sum_{m=0}^{M-1} A_{ms} \left( 1 - \frac{Y_{mk}}{A_{mk} F(\varphi, t_k) + \mu} \right)
\]
Using the notation defined in (13), (14), and (15), the PICD update can then be expressed as
\[
\hat{\varphi} \leftarrow \arg\min_{\varphi} \left\{ \Delta f(\hat{\varphi}_s, \varphi_s) \theta_1 + \frac{1}{2} \| \Delta f(\hat{\varphi}_s, \varphi_s) \|_{\tilde{\alpha}^2}^2 + \sum_{r \in \partial s} g_{s-r} \| T(\hat{\varphi}_s) - T(\varphi_r) \|_W^2 \right\}
\]
(30)
where
\[
\Delta f(\hat{\varphi}_s, \varphi_s) = \begin{bmatrix} \hat{a}_s, \hat{b}_s \end{bmatrix} \begin{bmatrix} \alpha(\hat{c}_s) & \beta(\hat{d}_s) \end{bmatrix} - \begin{bmatrix} a_s, b_s \end{bmatrix} \begin{bmatrix} \alpha(c_s) & \beta(d_s) \end{bmatrix}.
\]
(31)

We have found that the PICD update is best implemented using two-stage nested optimization.
\[
(c_s, d_s) \leftarrow \arg\min_{c_s, d_s \geq 0} \left\{ \Delta C([\hat{a}_s, \hat{b}_s, \hat{c}_s, \hat{d}_s], \varphi_s) \right\}.
\]
(32)

This nested optimization strategy is very important in reducing computation and assuring robust convergence. The inner optimization over \( \hat{a}_s \) and \( \hat{b}_s \) must be performed many times since this result is required for each update of outer optimization over \( \hat{c}_s \) and \( \hat{d}_s \). Fortunately, optimization over \( \hat{a}_s \) and \( \hat{b}_s \) can be done very efficiently with a simple steepest descent algorithm because this optimization does not require updating of \( \theta_1, \theta_2, \alpha(\hat{c}_s), \) or \( \beta(\hat{d}_s) \). Optimization with respect to \( \hat{c}_s \) and \( \hat{d}_s \) is done using iterative 1-D golden section search along the \( \hat{c}_s \) and \( \hat{c}_s + \hat{d}_s \) directions. This method assures the convergence is to a local minimum that meets the Kuhn-Tucker conditions [57]. Appendix A contains pseudocode that specify details of the algorithm.

**D. Multiresolution Initialization**

It is well known that for the tomographic problem the ICD reconstruction algorithm tends to have slow convergence at low spatial frequencies [58]. Normally, this problem is solved by initializing the ICD iterations with a FBP reconstruction. In this case, most of the residual error is only at high frequencies, so the ICD iterations converge quickly. However, for parametric reconstruction there is no simple direct reconstruction algorithm, such as FBP, to use as an initialization for the PICD iterations.

To solve this problem we use a multiresolution reconstruction scheme, which first computes coarse resolution reconstructions and then proceeds to finer scales. The coarsest resolution reconstruction is initialized with a single set of parameters obtained by weighted least squares curve fitting to the average emission rate of each time frame. Importantly, the average activity of each time frame can be calculated directly from the sinogram data with little computation. Finer resolution reconstructions are then initialized by interpolating the parametric reconstruction of the previous coarser resolution. This recursive process reduces computation because the computationally inexpensive reconstructions at coarse levels provide a good initialization for finer resolution reconstructions.

**IV. IMAGE DOMAIN PARAMETER ESTIMATION METHODS**

For purposes of comparison, we will also consider image domain methods which estimate parameters at each voxel from reconstructed images at each time. Each of these methods requires that the sinogram at each time frame be reconstructed using conventional reconstruction methods. For these methods, let \( x_s(t_k) \) denote the reconstructed activity of voxel \( s \) at time frame \( k \) collected at time \( t_k \), and let
\[
x_s = [x_s(t_0), x_s(t_1), \ldots, x_s(t_{K-1})]
\]
denote the activity of voxel \( s \) at all time frames.

**A. Pixel-wise Weighted Least Square (PWLS) Method**

The pixel-wise weighted least squares method estimates the parameters of each voxel by iteratively minimizing the weighted square error between the reconstructed time response of the voxel and the model output.

The parameters of voxel \( s \) are estimated as
\[
\hat{\varphi}_s = \arg\min_{\varphi_s} \| x_s - f(\varphi_s) \|_{W_s}^2
\]
(33)
where \( W_s \) is the \( K \times K \) diagonal weighting matrix for voxel \( s \). The weight of each time frame is chosen to be inversely proportional to the variance of the voxel activity in that time frame. This variance can be approximated by the activity estimate of this voxel, normalized by the duration of the time frame. In this case, \( W_s \) is a diagonal matrix with diagonal elements given by
\[
[W_s]_{k,k} = \frac{\Delta t_k}{\max\{x_{MIN,k}, x_s(t_k)\}}
\]
(34)
where \( \Delta t_k \) is the duration of time frame \( k \), and \( x_{MIN} \) controls the maximum allowable value for the weights.

The parameters are estimated using the same nested optimization strategy as specified in equation (32). In fact, this algorithm differs from the parametric reconstruction in only two respects. First, the data derivatives of equations (28) and (29) are replaced by
\[
\theta_1 = -2W_s (x_s - f(\varphi_s))
\]
(35)
\[
\theta_2 = 2W_s
\]
(36)
and second, the stabilizing functional \( S(\varphi) \) is set to 0.

**B. Pixel-wise Weighted Least Square Method with Spatial Regularization**

The spatial variation of the PWLS parameter estimates can be reduced by adding a stabilizing function to equation (33). The resulting estimate is given by
\[
\hat{\varphi} = \arg\min_{\varphi} \sum_{s=0}^{N-1} \| x_s - f(\varphi_s) \|_{W_s}^2 + S(\varphi)
\]
(37)
where $S(\cdot)$ is the spatial stabilizing functional [25], [26].

In the first method, which we call the pixel-wise least squares regularized (PWLSR) method, the stabilizing function has the form specified in equation (24). This is the same stabilizing function as was used for direct parametric reconstruction.

For the second method, which we call the PWLSZ method, we implemented the stabilizing functional described in [26]. This method smooths the PWLS estimate and uses it in the stabilizing function. Let $H(\cdot)$ be a smoothing operator and $\varphi^P$ be the PWLS parameter estimate. The constrained parametric image is then given by

$$\varphi^C = H(\varphi^P).$$

Next, a weight is calculated for each voxel. For voxel $s$ the corresponding weight is

$$w_s = \frac{\|x_s - f(\varphi^P)\|_2^2}{(\varphi^P_s - \varphi^C_s)^2}. $$

Using the constraint images and weights, the stabilizing function is given by

$$S(\varphi) = \sum_{s=0}^{N-1} w_s (\varphi_s - \varphi^C_s)^2. \quad (38)$$

Notice that the stabilizing function of (38) penalizes the difference between the parameters and a smoothed version of the parameters. Alternatively, the more traditional stabilizing function of (24) penalizes the spatial derivatives of the parameters.

For both of these methods the solution to (37) is computed using the nested optimization strategy specified in (32) and the data derivatives specified in (35) and (36).

C. Linear (Logan) Method

Kinetic parameter groups can sometimes be easily estimated by properly transforming the data. The Logan plot is a popular integral transform of the model given in equations (3), (4), and (8). This transformation can be expressed as follows.

$$\left[ \int_0^{t_k} \frac{x_s(t)dt}{x_s(t_k)} \right] = \frac{k_{1s}}{k_{2s}} \left( 1 + \frac{k_{3s}}{k_{4s}} \right) \left[ \int_0^{t_k} \frac{C_P(t)dt}{x_s(t_k)} \right] + \text{const}. \quad (39)$$

When the transformed variables (quantities in square brackets above) are plotted against each other, the resulting line has a slope equal to the compound parameter $VD_s$, which is defined in (2).

To calculate $BP_s$, the brain is segmented into a target region and a reference region. The target region consists of voxels within the brain that contain receptors for the tracer; and the reference region consists of the voxels that do not contain receptors for the tracer (i.e. $k_3 = 0$). Let $T$ be the set of voxel indices from target region, and $R$ be the set of voxel indices from reference region.

For a voxel $r \in R$ (from reference region), the distribution volume is

$$VD_r = \frac{k_{1r}}{k_{2r}}, \quad r \in R.$$

For each voxel $s \in T$ (from target region), the distribution volume ratio ($DV_{Rs}$) is

$$DV_{Rs} = \frac{VD_s}{\sum_{r \in R} VD_r} = 1 + \frac{k_{3s}}{k_{4s}}, \quad (40)$$

where $|R|$ denotes the number of voxels in the region $R$. Hence, the binding potential for the target region can be calculated as $BP_s = DV_{Rs} - 1$.

The assumptions that are used in the derivation of equations (39) and (40) are as follows:

- $k_1/k_2$ ratio is constant for every voxel in the brain (i.e., both target and reference regions)
- The tracer has high specific activity (so binding can be described as an apparent first order process)
- Blood volume fraction, $V_B$, is zero inside the target and the reference
- $k_3 = 0$ for all the voxels in the reference region

V. Computational Complexity Analysis

In order to better understand the computational requirements of parametric reconstruction, we derive expressions for the computational complexity of a number of parameter estimation algorithms.

First consider the PICD algorithm. For each voxel update, the data derivatives, $\theta_1$ and $\theta_2$, are computed once. The complexity of this computation is $O(KM_0)$, where $K$ is the number of time frames, and $M_0$ is the average number of nonzero projections per voxel. Assume the nested search described in equation (32) requires $L_{cd}$ evaluations of nonlinear parameters $\hat{c}_a$ and $\hat{d}_a$. Furthermore, assume that each update of $\hat{c}_a$ or $\hat{d}_a$ requires $L_{ab}$ evaluations of linear parameters $\hat{a}_s$ and $\hat{b}_s$.

Each evaluation with respect to $\hat{c}_a$ or $\hat{d}_a$ requires a convolution with the plasma input function and $L_{ab}$ evaluations with respect to $\hat{a}_s$ and $\hat{b}_s$. Let $K_c$ be the number of time points in the time-convolution kernel. Then the convolution requires $O(K_cK)$ operations, and the evaluation with respect to $\hat{a}_s$ and $\hat{b}_s$ requires $O(K)$ operations; so the total complexity of a voxel update is given by $O(KM_0 + L_{cd}(K_cK + L_{2ab}K))$, and the total complexity of PICD per full iteration for an $N$ voxel image is given by $O(KN(L_{cd}L_{ab} + K_cL_{cd} + M_0))$. The complexity of PWLS, PWLSR, and ICD, given in Table II, are then derived by removing the terms corresponding to operations that are not performed.

VI. Simulations

The following section compares the accuracy and computational burden of direct parametric reconstruction and image domain estimation methods.
<!DOCTYPE html>
<html><head></head><body><p>Parameter estimation from reconstruction

The blood contribution to the PET activity is assumed to be zero, and the tracer is assumed to have the form \( \lambda \). Total scan time is 60 min., divided into 18 time frames with 4 × 0.5 min, 4 × 2 min, and 10 × 5 min. The phantom had a resolution of 128 × 128 with each voxel having dimensions of (1.2 mm)³.

The rat phantom image at each time frame is forward projected into a sinogram using a Poisson model for the detected counts with a background (accidental coincidence) level of \( \mu = 0.001 \). Each sinogram consists of 180 angles and 200 radial bins per angle. This results in a value of \( M_0 \approx 934 \). A triangular point spread function with a 4 mm base width is used in forward projections. The blood function, \( C_P(t) \), is scaled so that the total number of counts in all sinogram frames is approximately 10 million.

### B. Algorithm Implementation

Direct reconstructions were computed using the PICD algorithm with three levels of multiresolution optimization corresponding to resolutions of 32 × 32, 64 × 64 and 128 × 128. The reconstructions used \( L_{cd} \approx 35 \) and \( L_{ab} = 15 \). In most cases, regularization was applied directly to the \( k_1, k_2, k_3, \) and \( k_4 \) parameters; so the stabilizing functional had the form

\[
S(k_1, k_2, k_3, k_4) = \sum_{i=1}^{4} \frac{1}{2 \sigma^2_{k_i}} \sum_{\{s,r\} \in \mathcal{N}} g_{s-r} |k_{i,s} - k_{i,r}|^2.
\]

(41)

### Table II

**Computational Complexity for a Single Full Iteration of PICD, PWLS, PWLSR, and ICD.** Notation: \( N \) = number of voxels; \( M_0 = 15 \) average number of projections per voxel; \( K = 15 \) the number of time frames; \( K_c = \) number of time points in the time-convolution kernel; \( L_{ab} = \) number iterations required for each update of \((a, b)\); \( L_{cd} = \) number iterations required for each update of \((c, d)\). Expressions do not include the computational cost of regularization.

### Table

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Function</th>
<th>Per Iteration Complexity</th>
</tr>
</thead>
<tbody>
<tr>
<td>PICD</td>
<td>Direct parametric reconstruction</td>
<td>( K N(L_{cd}L_{ab} + L_{cd} + M_0) )</td>
</tr>
<tr>
<td>PWLS</td>
<td>Parameter estimation from reconstruction</td>
<td>( K N(L_{cd}L_{ab} + K_cL_{cd}) )</td>
</tr>
<tr>
<td>PWLSR/PWLSZ</td>
<td>Regularized parameter estimation from reconstruction</td>
<td>( K N(L_{cd}L_{ab} + K_cL_{cd}) )</td>
</tr>
<tr>
<td>ICD</td>
<td>MAP image reconstruction</td>
<td>( K N(M_0) )</td>
</tr>
</tbody>
</table>

---

**Fig. 5.** Time-activity curves for 5 distinct tissue regions in rat brain phantom.

**Fig. 4.** Single-slice rat phantom. Regions of the rat phantom were derived from a segmented MR image. Different fill patterns indicate kinetically distinct tissue regions. Striatum is a region containing specific receptors for the tracer. Nonspecific-gray matter is tissue containing no specific binding sites for tracer but comparable blood flow parameters (k1, k2) to striatal area; cortex is modeled as containing low concentration of binding sites; white matter in our dynamic phantom contains no specific binding sites and low flow; non-brain, which comprises much of the slice has fast influx and efflux of tracer. Solid white areas in figure represent a mixture of background regions that do not contain any activity over time. The small white areas dorsal to (above) the striatum are ventricles that contain cerebral spinal fluid and no tracer. White areas surrounding brain correspond to skull which does not take up appreciable amounts of tracer.
where the function $g_{s-r}$ is inversely proportional to the distance between the voxels $s$ and $r$ and normalized to sum to 1, and the constants $\sigma_k^2$ control the regularization for each of the four parameters. The maximum likelihood (ML) estimate of $\sigma_k^2$ was computed for each parameter from the original parametric image using the formula [61]

$$\hat{\sigma}_k^2 = \frac{1}{N} \sum_{(s,r) \in N} g_{s-r} |k_i,s - k_{i,r}|^2 . \tag{42}$$

In the original formula, $N$ is the number of voxels in the image; however some parameter images have very few nonzero voxels, so we choose $N$ to be the number of nonzero voxels in the image. These ML parameters are then linearly scaled altogether to find a set of regularization parameters that minimize the RMSE of the estimated kinetic parameters. The resulting diagonal weighting matrix, $W$, from equation (24) has diagonal entries given by $W_{i,i} = \beta \frac{1}{\sigma_k^2(i)}$, where $\beta$ is the scaling factor that minimizes the parameter RMSE. Some results use regularization in the $k_1, k_2, BP, VD$ parameters, and $V D$ parameters. In this case, scaling parameters are selected similarly using the appropriate parameter values.

The image domain parameter estimation methods of section IV require that the image be reconstructed for each time frame. For this purpose, we used MAP image reconstruction with a quadratic prior (41) and a single fixed regularization parameter for all frame times. This single fixed parameter was chosen to minimize the total mean square error of the reconstructed emission image frames. The weighting matrix required for the PWLS, PWLSZ, and PWLSR algorithms was computed using (34) with $x_{MN} = 0.05$nCi/ml.

In order to compute the PWLSZ reconstruction as described in section IV-B, we smoothed the result of PWLS reconstruction with a $3 \times 3$ equal weight filter to calculate the constraints and weights. The weights were then scaled to minimize the MSE of the parameter estimates.

The PWLSR method was computed using a prior model on the $k_1, k_2, k_3,$ and $k_4$ parameters in a manner similar to that used for parametric reconstruction. As with parametric reconstruction, the $\sigma_k^2$ constants were first selected using the ML estimation method described above, and then scaled to yield the minimum RMSE estimates of the parameters.

For the linear (Logan) method, the cortex and striatum regions are selected as target regions, and the nonspecific-gray matter was used as the reference region. Since these regions were selected precisely from simulated data, all assumptions of this method are perfectly satisfied. A fixed number of iterations is used for each method. The multiresolution PICD method uses 30 iterations at $32 \times 32$ resolution, 20 iterations at $64 \times 64$ resolution, and 20 iterations at $128 \times 128$ resolution. Image domain methods use 15 iterations.

### C. Results

Figure 6 shows the reconstructions of the kinetic parameters. The first row contains the original parametric images. The remaining rows are respectively the reconstructions of PWLS, PWLSZ, PWLSR, PICD reconstruction regularized on $k_1, k_2, k_3,$ and $k_4,$ and PICD reconstruction regularized on $k_1, k_2, BP,$ and $VD$. In addition, the normalized RMSE of parameters $k_1, k_2, k_3,$ and $k_4$ estimated by these algorithms are listed in Fig. 7. The normalized RMSE of a parameter is computed as

$$\text{normalized RMSE}(k_i) = \sqrt{\frac{1}{|S|} \sum_{s \in S} (k_{i,s} - k_{i,s}^{estimated})^2} \cdot \sqrt{\frac{1}{|S|} \sum_{s \in S} (k_{i,s}^{estimated} - k_{i,s}^{PWLS})^2} ,$$

where $S$ is the domain where RMSE is computed, $|S|$ is the number of voxels in this domain, $k_{i,s}^{true}$ is the original value, $k_{i,s}^{PWLS}$ is the PWLS estimate of the parameter, and $k_{i,s}^{estimated}$ is the estimated value of the parameter for voxel $s$. The RMSE of $k_1$ is calculated over the whole image. The RMSE of parameters $k_2$ and $k_3$ are calculated over the support of $k_1$, and the RMSE of $k_4$ is calculated over the support of $k_3$.3

For the nonlinear parameters $k_3$ and $k_4$, the PWLS and PWLSZ methods both produced reconstructions which are very noisy, and this is reflected in the RMSE calculations. The PWLSR method with the GMRF prior produces lower RMSE reconstructions with more visually acceptable results for $k_3$ and $k_4$; however some details in these nonlinear parameters are lost. The parametric reconstruction regularized on $k_1, k_2, k_3,$ and $k_4$ produces higher SNR reconstructions than any of the image domain methods, and the reconstructed images are visually similar to the original phantom.

### TABLE III

<table>
<thead>
<tr>
<th>Region</th>
<th>$k_1$ min$^{-1}$</th>
<th>$k_2$ min$^{-1}$</th>
<th>$k_3$ min$^{-1}$</th>
<th>$k_4$ min$^{-1}$</th>
<th>$a$ min$^{-1}$</th>
<th>$b$ min$^{-1}$</th>
<th>$c$ min$^{-1}$</th>
<th>$d$ min$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Background</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>CSF</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nonspecific</td>
<td>0.0936</td>
<td>0.0968</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nonspecific-gray matter</td>
<td>0.0918</td>
<td>0.4484</td>
<td>1.2408</td>
<td>0.1363</td>
<td>0.0164</td>
<td>0.0706</td>
<td>1.7914</td>
<td>0.0312</td>
</tr>
<tr>
<td>Striatum</td>
<td>0.0918</td>
<td>0.4484</td>
<td>0.141</td>
<td>0.1363</td>
<td>0.0607</td>
<td>0.0311</td>
<td>0.628</td>
<td>0.09725</td>
</tr>
<tr>
<td>Cortex</td>
<td>0.0295</td>
<td>0.4484</td>
<td>0</td>
<td>0</td>
<td>0.02295</td>
<td>0</td>
<td>0.4484</td>
<td>0</td>
</tr>
</tbody>
</table>

KINETIC PARAMETERS USED IN THE SIMULATIONS FOR DISTINCT TISSUE REGIONS OF THE RAT HEAD.

3A very small amount of regularization was also used for $k_3$ and $k_4$ (i.e. $\sigma_{k_3}^2 = 1 \text{min}^{-2}$, $\sigma_{k_4}^2 = 0.1 \text{min}^{-2}$) to suppress impulsive noise in these reconstructions.

4When $k_1$ is zero, then $k_2$ and $k_3$ are not defined. Similarly, when $k_3$ is zero, $k_4$ is not defined.
However, the parametric reconstructions with regularization on \( k_1, k_2, BP, \) and \( VD \) yield the best quality results judging from both the visual quality and the computed RMSE.

For the comparison of parameters \( BP \) and \( VD \), spatial regularization is applied on \( k_1, k_2, BP, \) and \( VD \). In this case, the scaling of the four regularization constants are chosen to minimize the RMSE of the \( BP \) and \( VD \) estimates alone. The results are shown in Fig. 8 and the normalized RMSE of the estimates of all methods are given in Fig. 9. The RMSE of \( BP \) is estimated over the support of \( k_3 \), and the RMSE of \( VD \) is estimated over the support of \( k_1 \). Again, parametric image reconstruction produces the lowest RMSE estimation for both \( BP \) and \( VD \).

Once the parametric image is reconstructed, the ODE’s can be solved for any particular time to reconstruct the corresponding emission image. Fig. 10 compares these reconstructions to the conventional reconstructions computed using FBP and MAP reconstruction for time frames 5, 10, and 15. The FBP reconstructions use a Hamming filter with cutoff at the Nyquist frequency. The RMSE of these reconstructions for each frame and for total RMSE of all frames are given in Fig. 11.

Finally, the convergence speed as a function of CPU time for all algorithms is given in Fig. 12. The time needed to reconstruct emission images required by image domain methods is included in this figure. As can be seen from this

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**Fig. 6.** Parametric images of \( k_1, k_2, k_3 \) and \( k_4 \) estimated by the algorithms: (a) original (b) PWLS (c) PWLSZ (d) PWLSR (e) PICD1: PICD reconstruction (new method) regularized on \( k_1, k_2, k_3, \) and \( k_4 \) (f) PICD2: PICD reconstruction (new method) regularized on \( k_1, k_2, BP, \) and \( VD \).
figure, the convergence speed of direct parametric reconstruction is comparable to the pixel-wise methods. It has been shown that ICD has more rapid convergence at high spatial frequencies and relatively slower convergence at low spatial frequencies [58]. Therefore, we used multiresolution initialization to speed the convergence of lower frequency components in the parametric image. Table IV lists the CPU time required for a single iteration of each method. Notice that direct parametric reconstruction using PICD does not require substantially more computation per iteration than the image domain methods, and the image domain methods require that the images first be reconstructed. This result is consistent with the complexity listed in Table II since in this example, \((KcL_{cd} = 24, 185) >> (M_0 = 934) >> (L_{cd}L_{ab} = 525)\); so the computational complexity of the time convolution required for kinetic parameter estimation dominates the computations required for the tomographic reconstruction.

**VII. DISCUSSION**

In section VI-C, we demonstrated that the kinetic parameters estimated by the direct parametric image reconstruction (ie. the “direct”) have lower overall error as compared to those estimated in the image domain (i.e., the “indirect”).
Fig. 9. Normalized RMSE for the reconstructed $BP$ and $VD$. PICD reconstruction uses regularization on $k_1$, $k_2$, $BP$, and $VD$. Notice that PICD reconstruction gives the lowest RMSE results.

![Graph showing normalized RMSE for $BP$ and $VD$.]

Fig. 10. Activity images (a) original phantom (b) FBP reconstruction (c) MAP (d) PICD reconstruction (new method) for frames 5, 10, and 15.

![Activity images for different reconstruction methods.]

The improvement in the visual quality and the error of the kinetic parameter estimation may be due to the following factors:

- All the available data are used simultaneously.
- Kinetic parameters are estimated directly from PET sinogram data (for which we have a very good error model).
- Nonlinear estimation methods are used (so there is no need to linearize the model and introduce unwanted inaccuracy).
- Spatial regularization is done in the kinetic parameter domain (because neighboring voxels probably have similar function).

In contrast, the various image domain methods (described in Section IV) depend on the quality of the tomographic reconstructions of time-activity curves. Filtered backprojection (FBP) is still commonly used to reconstruct the dynamic PET data; unfortunately, it cannot produce the quality and the resolution achieved by the iterative reconstruction
techniques. Iterative methods (e.g., EM, ordered subset EM [62], or MAP [54]) require that images be reconstructed for each time frame and each slice. Therefore, direct parametric reconstruction reduces the dimensionality of the estimation problem from the number of time frames to the number of kinetic parameters in the compartment model. In our simulations, the dimensionality of the estimation problem was reduced by a factor of 4.5 (from 18 time frames to 4 parameters) by the direct method.

When using an image domain approach, spatial regularization can reduce the high spatial variance in the parametric images. We have found that spatial regularization based on a Gaussian Markov model produces less estimation error for all kinetic parameters except \( BP \), compared with a smoothing filter-based constraint.

The linear (Logan plot) method described in section IV-C is a very fast estimation technique. It tends to produce smooth images because it involves an integral transform of the data which suppresses noise. However, this method can only estimate some of the (compound) kinetic parameters. In receptor-ligand imaging, it provides no means for estimating \( k_1, k_2, k_3, \) or \( k_4 \), individually. Furthermore, to derive \( BP \) from distribution volume ratio, there must exist a reference region in the brain devoid of receptors (\( k_3 = 0 \)). For some tracers (e.g., muscarinic or nicotinic ligands), there is no readily apparent reference region and so the value of the Logan method is compromised. Even when an appropriate reference region exists in theory (e.g., for dopaminergic ligands) the validity of the parameter estimates in the rest of the tissue can be biased by the placement of (or spillover of activity into) the reference ROI. In our simulations, we use the precise target (striatum and cortex) and reference (nonspecific-gray matter) regions for this method which are selected from the original image.

Another drawback to linearizations of the model is that they achieve some of their computational simplicity by unmet model assumptions (e.g., that the blood volume fraction in the reference and target tissues is zero over all time). These simplifications have been shown to introduce biases that are aggravated with decreasing SNR [63]–[65]. Another common assumption that is implicit in the use of Logan-plot methods is that the \( k_1/k_2 \) ratio everywhere in the brain is constant (although we satisfy this constraint in our simulated data, the direct estimation method does not require it). This ratio can, of course, be regularized spatially in the direct method. Local regularization, however, is not nearly as rigid a requirement as expecting \( k_1/k_2 \) to be constant everywhere.

Although direct parameter estimation from the PET sinograms has been proposed previously as the EMPIRA algorithm [1], this or equivalent methods have not been fully implemented. This is likely due to the computational complexity of the M-step which was not fully specified, and the slow convergence of conventional EM iterations. With the development of computationally efficient and rapidly converging methods such as have been demonstrated in this paper, direct reconstruction to parametric images should become widely applied to dynamic PET data for which a kinetic model has been already established. It should be mentioned that there is nothing to prevent us from incorporating more complicated kinetic models into the PICD algorithm. Even though the solution to these models cannot be expressed in closed form, the power of our method, to decouple the (numerical) solution of the model from the other steps in the optimization procedure, is preserved.

We also believe that an extension of the PICD algorithm to list-mode data is possible. The function \( f(\phi_s) \), defined in equation (16), can be viewed as the coefficients of a zeroth order piece-wise constant spline. By using higher order splines, the activity of a voxel at any time can be computed. In this way, event arrival times can be incorporated into the probability function and log likelihood given in equations (19) and (20) respectively. In this case, \( f(\phi_s) \) would denote the spline coefficients for voxel \( s \), and the quantities \( \theta_1 \) and \( \theta_2 \) would be the first and second derivatives of the likelihood function with respect to the spline coefficients.

Our current implementation of direct reconstruction has certain limitations. For example, in image domain estimation methods it is possible to register the images for motion compensation. External measurement devices can allow us to record motion during acquisition [66] and correct the data in an automated fashion. The current implementation of our algorithm does not allow for this type of compensation. Another limitation is that our method assumes that all voxels are well modeled by the same family of model kinetics, which might not be the case in practice. However we note that single families of model kinetics have been sufficient for describing receptor ligands in different regions of the brain on an ROI-basis.
VIII. Conclusions

In this paper, we introduce a method for the direct reconstruction of kinetic parameters at each voxel from dynamic PET sinogram data. Our algorithm, which we call parametric iterative coordinate decent (PICD), decouples the nonlinearities between the tomographic model, the kinetic model, and the regularized parameters. It also allows one to regularize with respect to any desired parametrization, even if the parameters that are selected are nonlinearly related to the projections or the kinetic model parameters. Using an anatomically and physiologically realistic small animal phantom, we demonstrated that our method can reduce the mean squared error in model parameter estimates; and we show that for our example, it does not require substantially more computation than more conventional methods for computing dense parameter estimates in the image domain.

Acknowledgement

Authors would like to thank Cristian Constantinescu, Chunzhi Wang, Dr. Karmen Yoder and Dr. Ti-Qiang Li at Indiana University School of Medicine for their help in constructing the rat phantom from MR data.

A. Appendix - Pseudocode

\[ \varphi \leftarrow \text{ParametricReconstruct(sinograms)} \]

\{ for each iteration \}

\{ for each voxel \}

\[ [\theta_1, \theta_2] \leftarrow \text{ComputeDerivatives(sinograms, } \varphi]\]

\[ \hat{\varphi}_s \leftarrow \varphi_s \]

\[ [\tilde{\alpha}, \tilde{\beta}] \leftarrow \text{ConvolveWithPlasma}(\hat{\varphi}_s, \hat{d}_s, [t_0, \ldots, t_{K-1}], \lambda, V_B, C_F) \]

\[ \tilde{c}_s \leftarrow \arg \min_{\tilde{c}_s} \{ \text{CostFunction}([\tilde{a}_s, \tilde{b}_s, \tilde{c}_s, \tilde{d}_s], \varphi_s, \alpha, \beta, \theta_1, \theta_2; \{ \varphi_r : r \in \partial S \}, W) \} \]

\[ [\tilde{a}_s, \tilde{b}_s] \leftarrow \text{EstimateAandB} \left( \tilde{\varphi}_s, \varphi_s, \tilde{\alpha}, \tilde{\beta}, \alpha, \beta, \theta_1, \theta_2, \{ \varphi_r : r \in \partial S \}, W \right) \]

\[ l \leftarrow \arg \min_l \{ \text{CostFunction}([\tilde{a}_s, \tilde{b}_s, \tilde{c}_s + l, \tilde{d}_s + l], \varphi_s, \alpha, \beta, \theta_1, \theta_2; \{ \varphi_r : r \in \partial S \}, W) \} \]

\[ \tilde{c}_s \leftarrow \tilde{c}_s + l; \quad \tilde{d}_s \leftarrow \tilde{d}_s + l \]

\[ [\tilde{a}_s, \tilde{b}_s] \leftarrow \text{EstimateAandB} \left( \tilde{\varphi}_s, \varphi_s, \tilde{\alpha}, \tilde{\beta}, \alpha, \beta, \theta_1, \theta_2, \{ \varphi_r : r \in \partial S \}, W \right) \]

\[ \hat{\varphi}_s \leftarrow \tilde{\varphi}_s \]

\}

\}

\[ \text{cost} \leftarrow \text{CostFunction}(\tilde{\varphi}_s, \varphi_s, \alpha, \beta, \theta_1, \theta_2; \{ \varphi_r : r \in \partial S \}, W) \]

\{ \[ [\tilde{a}_s, \tilde{b}_s] \leftarrow \text{ConvolveWithPlasma}(\hat{\varphi}_s, \hat{d}_s, [t_0, \ldots, t_{K-1}], \lambda, V_B, C_F) \]

\[ [\tilde{a}_s, \tilde{b}_s] \leftarrow \text{EstimateAandB} \left( \tilde{\varphi}_s, \varphi_s, \tilde{\alpha}, \tilde{\beta}, \alpha, \beta, \theta_1, \theta_2, \{ \varphi_r : r \in \partial S \}, W \right) \]

\[ \text{cost} \leftarrow \text{DeltaCost}(\tilde{\varphi}_s, \varphi_s, \tilde{\alpha}, \tilde{\beta}, \alpha, \beta, \theta_1, \theta_2; \{ \varphi_r : r \in \partial S \}, W) \}

\}

\begin{align*}
\text{for } L_{\text{ab}}/3 \text{ iterations } \}

\Delta C & \leftarrow \text{DeltaCost}([\tilde{a}_s, \tilde{b}_s, \tilde{c}_s, \tilde{d}_s], \varphi_s, \tilde{\alpha}, \tilde{\beta}, \alpha, \beta, \theta_1, \theta_2, \{ \varphi_r : r \in \partial S \}, W) \}

\Delta C & \leftarrow \Delta f \theta_1 + \frac{1}{2} \| \Delta f \|_2 + \sum_{r \in \partial S} g_{s,r} \| T(\hat{\varphi}_s) - T(\varphi_r) \|_W \}

\end{align*}

References


